The enlargement of either the pulmonary artery or the aorta can result in a left-sided recurrent nerve palsy, although this has rarely been reported as a complication of giant cell arteritis (5). Cardiac involvement is also rare in giant cell arteritis, and it usually appears as coronary arteritis and myocardial infarction (6). Pulmonary involvement is a rare complication and can be the cause of respiratory symptoms. Its diagnosis is challenging, requiring often the use of positron emission tomography/CT to assess the exact extension of arteritis (7). Unfortunately, this method was unavailable at that time in our center.

The gold standard of the diagnosis is temporal artery biopsy, although this can give a false-negative result in 15%–40% of the cases (8). Because of a great possibility that the temporal artery was not affected, temporal artery biopsy was considered unnecessary.

In the present case, the diagnostic difficulties were caused by the atypical appearance of giant cell arteritis. The vascular involvement was specific to both Takayasu- and giant cell arteritis, although our patient was much older than 50 years at the onset of the disease, which would have been very uncommon in Takayasu arteritis. In addition, some of the main manifestations of temporal arteritis were also missing. Although the autoimmune serology suggested the presence of granulomatosis with polyangiitis, the clinical appearance was against small vessel vasculitis. The ophthalmologic examination with electrophysiological examination led to the final proof: According to the literature, arteritic anterior ischemic optic neuropathy is associated with giant cell arteritis in most of the cases (2).

**Conclusion**

Pericardial effusion is still an extraordinary manifestation of temporal arteritis, but the exact prevalence is unknown. Although it is usually asymptomatic, it can lead to the final diagnosis, and the development of aortic aneurysm can be prevented.

**Acknowledgments:** The corresponding author gratefully acknowledges the teaching efforts of the co-authors and the assistance of the nurses in the 2nd Department of Internal Medicine of University of Szeged.

**Informed consent:** Written informed consent was obtained from the patient.

**Video 1.** Thickened aortic wall on transesophageal echocardiography: The ascending aortic wall was 9 mm and the aortic arch was 5 mm thick.

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DOI:10.14744/AnatolJCardiol.2019.00502

**Happy heart syndrome mimicking acute pulmonary embolism and acute coronary syndrome 🦋**

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**Introduction**

Takotsubo cardiomyopathy (TC), also named as the “broken heart syndrome (BHS),” “apical ballooning syndrome,” or “stress-induced cardiomyopathy,” is characterized by transient left ventricular systolic dysfunction without an evidence of obstructive coronary artery disease (1). Postmenopausal women account for approximately 90% of the cases (1). Recent studies have shown that TC can be triggered by negative as well as positive life events, owing to which it is also known as the “happy heart syndrome” (HHS) (2). However, it is frequently triggered by physically or emotionally stressful events such as the receipt of bad news; incidents of death, accidents, and natural disasters; or presence of complicated medical diseases as well as events...
including surgical procedures, sepsis, trauma, or cerebrovascular diseases (3). Although acute pulmonary embolism has been listed as a potential contributor in TC, there are currently no case reports of TC mimicking acute pulmonary embolism in the literature. Here, we present a rare case of HHS mimicking acute pulmonary embolism that completely recovered with anticoagulant and diuretic therapies.

Case Report

A 65-year-old woman with no cardiovascular risk factors presented to our emergency department (ED) with retrosternal chest pain at rest. Her medical history included only hypothyroidism, which was being treated with 50 μg levothyroxine sodium once daily. The patient was admitted to the ED with a blood pressure of 84/45 mm Hg and a heart rate of 98 bpm. She was afebrile, with a respiratory rate of 25 breaths/min and an oxygen saturation of 95%. Physical examination revealed no jugular venous distention or lower extremity edema, and the lungs were clear on auscultation. The ECG showed sinus rhythm and an S1Q3T3 pattern (Fig. 1).

The patient’s complete blood count and renal and liver function tests were all within the normal ranges. The routine laboratory tests conducted in ED revealed that the level of high-sensitivity cardiac troponin T was 86 ng/L (normal reference range, 0–52 ng/L), that of creatine kinase-MB was 8.61 ng/mL (normal reference range, 0–4.99 ng/mL), and that of D-dimer was 1213 ng/mL (normal reference range, 0–500 ng/mL). Echocardiogram was performed upon admission to ED, and the results showed normal tricuspid annular plane systolic excursion and a right ventricular diameter of 18 mm, an enlarged left ventricle of 51 mm at end-diastole with a reduced left ventricular ejection fraction of 32%, and a decreased motion of the left ventricular anterior–anteroseptal wall and apex (Fig. 2a, Video 1). Acute myocardial infarction was suspected due to regional wall motion abnormality and reduced left ventricular systolic function. On the other hand, subsegmental acute pulmonary embolism was also suspected due to the S1Q3T3 pattern on ECG and the slightly increased D-dimer levels. The patient was administered 325 mg of aspirin, 600 mg oral loading dose of clopidogrel, enoxaparin (30 mg intravenous bolus and 1 mg/kg twice a day subcutaneously 15 min after bolus), and 40 mg of intravenous furosemide in ED. The patient was transferred to the catheterization laboratory within 30 min of the first medical contact. However, coronary angiography revealed normal coronary arteries. Furthermore, computed tomography with angiogram was performed to rule out pulmonary embolism, which revealed normal pulmonary arteries without any thrombi. The detailed history obtained retrospectively revealed that the patient’s complaints started immediately after learning that she would be a grandmother. Her hospital course was uneventful and her pain resolved within 3 h of admission. She was treated with standard heart failure therapy including angiotensin-converting enzyme inhibitors, beta-blockers, and diuretics.

Repeat echocardiography performed on day 1 displayed a normal-sized left ventricle with left ventricular apical hypokinesia and a left ventricular ejection fraction of 48%. On day 3, transthoracic echocardiogram revealed normal left ventricular systolic functions without any segmental wall motion abnormality (Fig. 2b). The ejection fraction was 64%. The patient was diagnosed with HHS and she was discharged home with routine patient–clinic appointment.

Discussion

In 1991, Dote et al. (4) first described TC in a series of five case in a Japanese population. In this syndrome, which was named after the shape of the octopus trap (takotsubo), the heart appears narrowed at the base with apical ballooning. Although the pathogenesis of HHS and BHS is not well understood, several etiologies have been proposed including catecholamine excess, coronary artery vasospasm, and estrogen deficiency (1-4). Although this syndrome has been named BHS, in 2016, Ghadri et al. (2) demonstrated that pleasant events, such as a surprise party, positive job interview, and winning jackpots, may trigger TC. The authors described 20 patients with this unique manifestation and named it HHS. Clinical presentation of patients with HHS was similar to those with the BHS, including symptoms, electrocardiographic and laboratory findings, and long-term outcomes (2).
Because TC shares similar clinical presentations with acute coronary syndrome and acute pulmonary embolism, it is easily misdiagnosed. Although acute pulmonary embolism is reported as a trigger for TC, the concurrence of TC and pulmonary embolism has rarely been reported (5).

**Conclusion**

The present patient was diagnosed with HHS based on the clinical, laboratory, angiographic, electrocardiographic, and echocardiographic findings. In cases that present with overlapping symptoms, it may be difficult to distinguish TC from pulmonary embolism and acute coronary syndrome. A history of recent emotional trauma, whether it was triggered by bad or good news, can prompt a clinician to consider TC as a diagnosis. The present case illustrates the challenges of distinguishing TC from acute coronary syndrome or pulmonary embolism. It is crucial that ED physicians recognize BHS and HHS as rare yet considerable reasons of acute chest pain and dyspnea. An assessment of the patient’s medical history, particularly of previous emotional or physical triggers, is fundamental in the diagnosis of TC.

**Informed consent:** Written informed consent was obtained from the patient.

**Video 1.** Apical four-chamber view (left side of the video) in transthoracic echocardiography performed on admission showing akinesis. Apical four-chamber view (right side of the video) in transthoracic echocardiography performed on day 3 showing the complete recovery of left ventricular systolic function.

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DOI:10.14744/AnatolJCardiol.2019.04052